## **Raw Sequence Listing Error Summary**

	ERROR DETECTED	SUGGESTED	CORRECTION	SERIAL NUMBER: $\underline{\mathcal{O}}$	<u>9/189,41</u> 5
ATTN	: NEW RULES CASES: P	LEASE DISREGA	ARD ENGLISH "ALPI	HA" HEADERS, WHICH WERE INSERTED	BY PTO SOFTWARE
1	Wrapped Nucleics	The number/text	at the end of each lin	e "wrapped" down to the next line.	
	•••	This may occur i	f your file was retrieve	ed in a word processor after creating it.	
	Please adjust your right margin to .3, as this will prevent "wrapping".				
2	Wrapped Aminos	The amino acid r	number/text at the end	f of each line "wrapped " down to the next li	ne.
	This may occur if your file was retrieved in a word processor after o			. • •	
		•		as this will prevent "wrapping".	
3	Incorrect Line Length	The rules require	that a line not excee	d 72 characters in length. This includes spa	aces.
4	Misaligned Amino Acid	The numbering u	nder each 5th amino	acid is misaligned. This may be caused by	the use of tabs
	Numbering	between the numbering. It is recommended to delete any tabs and use spacing between the numbers.			
5	Non-ASCII	This file was not	saved in ASCII (DOS	) text, as required by the Sequence Rules.	
		Please ensure yo	our subsequent submi	ssion is saved in ASCII text so that it can be	e processed.
6	Variable Length			which represented more than one residue.	
		-		ly represent a single residue.	
		Please present the	ne maximum number	of each residue having variable length and	_
		indicate in the (ix	() feature section that	some may be missing.	g from amino acid s section from the >-<223> section
7	Patentin ver. 2.0 "bug"	A "bug" in Patent	In version 2.0 has car	used the <220>-<223> section to be missing	from amino acid
	•			, Patentin would automatically generate this	s section from the 😞 📆
				ce. Please manually copy the relevant <220	)>-<223> section
			nt amino acid sequen		J T
					1631
8	Skipped Sequences	Sequence(s)	missing If intention	al, please use the following format for each	skinned sequences
	(OLD RULES)		ON FOR SEQ ID NO:		0
	(OLD NOLLO)			v. (Do not insert any headings under "SEQUE	NCE CHARACTERISTICS")
			DESCRIPTION:SEQ		TOE OFFICIOUS ,
		• •			
		inis sequence i	s intentionally skipp	ea	
		Please also adjus	st the "(iii) NUMBER (	OF SEQUENCES:" response to include the	skipped sequence(s).
9	Skipped Sequences	Sequence(s)	_ missing. If intention	al, please use the following format for each	skipped sequence.
	(NEW RULES)	<210> sequence	e id number	•	
١		<400> sequence	e id number		
1		000			
, J	Use of n's or Xaa's	Lise of n's and/or	Xaa's have been det	ected in the Sequence Listing.	
	(NEW RULES)			Y if n's or Xaa's are present.	
	(NEW NOCES)			lain location of n or Xaa, and which residue	n or Xaa represents.
1	Use of <213>Organism	Sequence(s)	are missing this	mandatory field or its response.	
'	(NEW RULES)	Ocquence(3)	are missing this	inalidatory field of his response.	
2	Use of <220>Feature	Sequenco(s)	are missing the <20	0>Feature and associated headings.	•
			- •	<del>-</del>	nown"
	(NEW RULES)			Y if <213>ORGANISM is "Artificial" or "Unk	HOWII
		•	-	terial in <220> to <223> section.	
		(See "Federal	Register," 6/01/9	8, Vol. 63, No. 104, pp. 29631-32)	(Sec. 1.823 of new Rules)
3	Patentin ver. 2.0 "bug"			nction of Patentin version 2.0. This cause	
		file, resulting in m	issing mandatory nur	neric identifiers and responses (as indicated	I on raw sequence listing).
	2. A	Inches de minera de	PCilo Managed of	now other means to convilie to floorly disk	

AKS-Biotechnology Systems Branch- 5/15/99